# **PCT**

# WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



## INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 6:		(11) International Publication Number: WO 95/32790
B01D 53/94, B01J 23/38, 23/89	A1	(43) International Publication Date: 7 December 1995 (07.12.95)
(21) International Application Number: PCT/USS		ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE).
(22) International Filing Date: 31 May 1995 (3	31.05.9	
(30) Priority Data: 08/252,100 1 June 1994 (01.06.94)	τ	Published  With international search report.  Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.
(71) Applicant: ALLIEDSIGNAL ENVIRONMENTAL LYSTS [US/US]; Alliedsignal Inc., 101 Colomb P.O. Box 2245, Morristown, NJ 07962-2245 (US).	ia Roa	
(72) Inventors: KHARAS, Karl, C., C.; 7766 South Fulto Tulsa, OK 74136 (US). ROBOTA, Heinz, Juerge South Erie Avenue, Tulsa, OK 74137 (US). NUNA Gerard; 8729 South 68th East Avenue, Tulsa, O. (US). HENK, Michael, G.; 1220 East 18th Street OK 74120 (US).	n; 112 AN, Joh K 741	0 n, 3
(74) Agent: CRISS, Roger, H.; AlliedSignal Inc., Law Deg McNally), 101 Columbia Road, P.O. Box 2245, Mo NJ 07962-2245 (US).	pt. (C. orristow	h.
·		

(54) Title: ALLOYED METAL CATALYSTS FOR THE REDUCTION OF NO<sub>x</sub> IN THE EXHAUST GASES FROM INTERNAL COMBUSTION ENGINES CONTAINING EXCESS OXYGEN

#### (57) Abstract

Removal of carbon monoxide, hydrocarbons, and nitrogen oxides from the exhaust gas from lean-burn, diesel and other engines which produce exhaust gases containing oxygen in substantial excess of stoichiometric is provided by catalysts containing platinum alloyed with cobalt, nickel, copper, rhodium, palladium, silver or gold. Alloying is accomplished by thermal treatment or by colloidal methods. The catalysts are capable of reducing nitrogen oxides (NO<sub>x</sub>) within a predetermined range of engine operating temperatures.

# FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AT	Austria	GB	United Kingdom	MR	Mauritania
ΑU	Australia	GE	Georgia	MW	Malawi
BB	Barbados	GN	Guinea	NE	Niger
BE.	Belgium	GR	Greece	NL	Netherlands
BF	Burkina Faso	HU	Hungary	NO	Norway
BG	Bulgaria	IE	Ireland	NZ	New Zealand
BJ	Benin	IT	Italy	PL	Poland
BR	Brazil	JP	Japan	PT	Portugal
BY	Belarus	KE	Кепуа	RO	Romania
CA	Canada	KG	Kyrgystan	RU	Russian Federation
CF	Central African Republic	KР	Democratic People's Republic	SD	Sudan
CG	Congo		of Korea	SE	Sweden
CH	Switzerland	KR	Republic of Korea	SI	Slovenia
CI	Côte d'Ivoire	KZ	Kazakhstan	SK	Slovakia
CM	Cameroon	LI	Liechtenstein	SN	Senegal
CN	China	LK	Sri Lanka	TD	Chad
CS	Czechoslovakia	LU	Luxembourg	TG	Togo
CZ	Czech Republic	LV	Larvia	TJ	Tajikistan
DE	-	MC	Monaco	TT	Trinidad and Tobago
DK DK	Germany Denmark	MD	Republic of Moldova	UA	Ukraine
	= = = = = = = = = = = = = = = = = = = =	MG	Madagascar	US	United States of America
ES	Spain	ML	Mali	UZ	Uzbekistan
FI	Finland	MN	Mongolia	VN	Viet Nam
FR	France	WIIA	MOIBOIR	***	· · · · · · · · · · · · · · · · ·
GA	Gahon				

10

15

20

25

30

35

# ALLOYED METAL CATALYSTS FOR THE REDUCTION OF NO<sub>X</sub> IN THE EXHAUST GASES FROM INTERNAL COMBUSTION ENGINES CONTAINING EXCESS OXYGEN

BACKGROUND OF THE INVENTION

The invention relates generally to the control of hydrocarbons (HC), carbon monoxide (CO), and nitrogen oxides  $(NO_X)$  in the exhaust of internal combustion engines. More particularly, the invention relates to the removal of  $NO_X$  when the exhaust gases include oxygen substantially in excess of that needed for combustion of the fuel. This is typically the case with leanburn engines, diesel engines, and other engines currently under development.

In recent years three-way catalysts have been used to remove all of the three principal noxious components in auto exhaust gases. The engines are run with stoichiometric air/fuel ratios and the catalysts are able to remove all three components at the same time, that is, a single catalyst is sufficient over the range of engine operating temperatures. More recently, development of socalled "lean-burn" engines is being driven by the desire to improve fuel economy. However, such engines operate with air-fuel ratios which are far from the typical stoichiometric conditions. Instead of an air-fuel ratio of about 14.55/1 by weight, the lean-burn engine may operate with air-fuel ratios above 18/1, up to about 22-24/1, or even higher ratios of 28-30/1 for advanced lean-burn engines or diesel engines. Under such conditions the engine exhaust will include more hydrocarbons, less carbon monoxide, and less, but still excessive nitrogen oxides. While an oxidation catalyst is capable of removing hydrocarbons and carbon monoxide, since the oxygen content is high, say about 3-10% by volume, it is clear that conditions are not favorable for the reduction of nitrogen oxides. Conventional gasoline engines that operate near the stoichiometric point can experience lean transients where the air-fuel ratio reaches about 15/1. Under these conditions, the chemical equivalence ratio, defined as the sum of oxidizing chemical equivalents divided by the sum of reducing chemical equivalents, of emissions from the engine can reach a value of about 2. (At the stoichiometric point the equivalence ratio is 1.0.) The present inventors are concerned with nitrogen oxide reduction from lean-burn and diesel engines where the chemical equivalence ratio is greater than 3, often greater than 5, and frequently greater than 10.

10

15

25

30

35

There are two generally recognized routes to removing nitrogen oxides. First, the nitrogen oxides can be decomposed to the diatomic nitrogen and diatomic oxygen. This reaction is thermodynamically favored, but catalysts which are able to carry out this reaction under the highly oxidizing conditions and high temperatures found in engine exhaust have not yet been found. The second route is the chemical reduction of nitrogen oxides using reducing agents present in the exhaust, such as carbon monoxide, hydrocarbons, and hydrogen. This is considered to be the mechanism of the three-way catalyst. However, such catalysts see the exhaust from an engine operating with a stoichiometric airfuel ratio which contains little oxygen. When a large excess of oxygen is present, as in a lean-burn or diesel engine, the oxygen, being much greater in volume than the nitrogen oxides would be expected to react with the hydrocarbons, carbon monoxide, and hydrogen, thus removing the reducing agents needed to reduce nitrogen oxides.

In commonly-assigned USSN 07/990,216 a sequence of catalysts was used to cover the range of operating temperatures experienced with internal combustion engines. The second and third catalysts in the sequence were examples of catalysts of the present invention.

#### 20 SUMMARY OF THE INVENTION

In one aspect, the invention is a catalyst which is capable of removing carbon monoxide, hydrocarbons, and nitrogen oxides from the exhaust gases produced by an engine which is operated with a substantial excess of air relative to that required for stoichiometric combustion of the fuel, such as leanburn, diesel, and the like. The catalysts of the invention employ alloys of platinum with cobalt, nickel, copper, rhodium, palladium, silver, or gold dispersed on an inert support. The preferred alloys are platinum and rhodium, platinum and silver, platinum and cobalt, and platinum and palladium. Ternary alloys containing platinum, rhodium, and a third metal chosen from the group consisting of cobalt, palladium and nickel also are useful. The atomic ratios of platinum to silver may vary from about 1/10 to 1000/1. The atomic ratios of platinum to rhodium or palladium may vary from about 1/2 to 1000/1. The atomic ratio of platinum to cobalt, nickel, or copper may vary from about 1/1 to about 1000/1. The atomic ratio of platinum to gold may vary from about 15/1 to about 1000/1.

15

20

25

30

Base metals may be included as stabilizers or promoters, provided that they do not adversely affect the alloying of platinum and the other metals.

The alloying process may involve treatment of an impregnated support in an inert atmosphere containing up to about 30% steam and up to 100% hydrogen at temperatures of about 600° to 1200°C. Oxygen is typically excluded but may be present when hydrogen is absent and when it does not interfere with the desired metal alloying. Supports stable under such conditions include barium-stabilized alumina,  $\alpha$ -alumina,  $\delta$ -alumina,  $\theta$ -alumina, zirconia, titania, silicon carbide, and cordierite.

When the metal to be alloyed with platinum is cobalt, nickel, copper, palladium, rhodium, silver, or gold on a temperature stable support, the metals, after being deposited on the support, are exposed to an inert atmosphere containing 0-30 vol.% steam, greater than zero up to 100 vol.% hydrogen, and zero up to less than 100 vol.% inert gas at temperatures in the range of about 600° to 1200°C.

When the metal to be alloyed with platinum is palladium, rhodium, or gold, hydrogen may be omitted and up to 21 vol.% oxygen included. The temperature is selected so that the metal to be alloyed with platinum is favored rather than the metal oxide, for example, a temperature at which Rh<sub>2</sub>O<sub>3</sub> is unstable with respect to rhodium metal.

Alloyed catalysts also may be prepared by applying colloidal suspensions of alloys to a support, which may be selected from the stable supports mentioned above or alternatively other less stable supports including γ-alumina, crystalline boehmite, crystalline alumina silicates, non-crystalline silica-alumina, α-Zr(HPO4)2, NaZr<sub>2</sub>(PO<sub>4</sub>)3, and (Na,H)Zr<sub>2</sub>(PO<sub>4</sub>)3. In this process, the alloys are formed in solution or in suspension and applied to the support by adsorption; the high temperature treatment described above is not required.

# Brief Description of the Drawing

Figure 1 is a plot of NO conversion versus temperature for catalysts tested in Example 3.

Figure 2 is a plot of NO conversion versus temperature comparing the alloyed and unalloyed catalysts of Example 4.

Figure 3 shows the x-ray diffraction pattern of an alloyed Pt-Rh catalyst.

15

20

25

30

35

#### Description of the Preferred Embodiments

Those working in this field intend to either decompose nitrogen oxides into the elements directly or to reduce them selectively using reducing agents under oxidizing conditions. The catalysts and catalyst systems of the present invention are considered to function by reducing the nitrogen oxides rather than decomposing them. However, that conclusion was reached based on experimental evidence and is not an essential aspect of the invention. The reaction mechanisms by which nitrogen oxides are reduced are believed to vary depending on the catalyst and the operating temperatures.

It has been discovered that catalysts may function differently depending upon the operating conditions and that each catalyst has a temperature range within which it is capable of destroying nitrogen oxides. We will refer to this temperature range as a "window" within which nitrogen oxides are reduced. This is generally contrary to the experience with three-way catalysts which become active ("light-off") at a temperature of about 200° to 350 °C and thereafter are able to oxidize hydrocarbons and carbon monoxide while reducing nitrogen oxides over the usual range of operating temperatures, say about 300° to 800°C. It should be noted here that auto exhaust composition and temperature vary as driving conditions change. Consequently, tests of auto exhaust catalysts require that a car be operated over a range of conditions representing typical driving. Once three-way catalysts have reached operating temperature the performance is not greatly affected by the exhaust gas temperature. This is not the case with catalysts for engines operating in the leanburn mode, where catalysts have a temperature range within which they are effective to remove nitrogen oxides, but not outside that range. As will be appreciated, such a characteristic is not compatible with the usual variation in engine exhaust temperatures. However, by proper selection of catalysts and their pretreatment, it is possible to remove nitrogen oxides from temperatures of about 200° to 700°C.

Selection of catalysts for this difficult task requires consideration of the characteristics of each catalyst. It is believed that the chemical reactions differ with the catalyst composition and the temperature of operation. Typically, a catalyst has a temperature range within which it is effective and outside that range the conversion of nitrogen oxides is minimal. At the same time it is still necessary to remove the reducing agents, such as residual hydrocarbons, hydrogen, and carbon monoxide. Generally, workers in the field have suggested

15

20

25

30

using more than one catalyst, for example, a catalyst to convert nitrogen oxides followed by an oxidation or three-way catalyst. Based on the results of our experiments we believe that certain catalysts may be capable of reducing nitrogen oxides within a particular temperature range but still remain very effective for oxidizing the reducing agents i.e., hydrocarbons and carbon monoxide, at other temperatures. Thus, it appears that providing for both the reduction of nitrogen oxides and the oxidation of the reducing agents involves not only selection of the proper catalysts but the proper combination of catalysts and their positioning in order to achieve the desired reduction of nitrogen oxides throughout the range of operating conditions of a lean burn engine.

#### Catalyst Supports

As shown in USSN 07/990,216, a sequence of catalysts may be employed to cover the expected range of operating temperatures. The second and third catalysts of the preferred embodiment are alloyed noble metal catalysts of the present invention.

The second catalyst employs a new support, which is a barium stabilized delta alumina. It combines noble metals, particularly platinum and rhodium with cobalt in an alloyed form, which provide advantages relative to the comparable non-alloyed metals.

The support for the second catalyst is prepared by a sol-gel procedure which produces a very stable delta alumina and, although it contains a substantial amount of barium, appears to be solely alumina when examined by x-ray diffraction. This support designated BaO-δ-Al<sub>2</sub>O<sub>3</sub> is the subject of USSN 07/994,838 incorporated herein by reference. In one example, the process for preparing the BaO-δ-Al<sub>2</sub>O<sub>3</sub> support employs the following steps.

- (1) an aqueous barium acetate solution is prepared having a concentration suitable to provide about 1 to 12 wt.% Ba in the product;
- (2) an aqueous boehmite sol having particles above about 150 Å (15 nm), preferably below 300 Å (30 nm) is stirred;
- (3) the stirred sol of (2) is combined with the aqueous barium acetate solution of (1) and about 2 to 30 wt.% (preferably 10 to 25 wt.%) of polyethylene glycol (PEG) relative to the sum of PEG plus water to form a gel;
- 35 (4) the gel produced in step (3) is dried and then calcined in air to 1000°C;

10

15

(5) the calcined gel of (4) is then treated at about 1000° to 1350°C, preferably about 1140°C, in an atmosphere containing about 0 to 30 % steam, preferably about 10% steam, to convert the alumina into an essentially pure delta alumina.

This unique delta alumina is very stable and at temperatures up to 1350°C has been found to retain the delta form with only a small amount of theta alumina and no alpha alumina being formed.

Another very stable catalyst support is alpha alumina. This material is not usually associated with highly active catalysts since it has a low surface area. Other forms of alumina having greater surface area, e.g., gamma, theta, and delta aluminas are usually used. However, for catalysts of the invention, alpha alumina is a useful support.

Other temperature-stable supports for use when thermal processing is used to form alloys include  $\delta$ -alumina,  $\theta$ -alumina, zirconia, titania, silicon carbide, and cordierite. When alloys are preformed and applied as colloidal solutions, the temperature-stable supports just mentioned may be used, but other less stable supports also may be chosen, including  $\gamma$ -alumina, crystalline boehmite, crystalline alumina silicates, non-crystalline silica-alumina,  $\alpha$ -Zr(HPO4)2, NaZr<sub>2</sub>(PO4)3, and (Na,H)Zr<sub>2</sub>(PO4)3.

20

25

30

35

### Alloved Noble Metals

The catalysts of the invention are alloys of platinum with other metals and include alloys of platinum with elements of Group 9, 10, and 11 of the IUPAC Periodic Table. Heretofore, it has been thought that alloying was detrimental to catalyst performance. See, for example, J.T. Kummer, J. Phys. Chem. 1986, 90, 4747-4752. Also, sintering of the noble metal crystallites has been seen in aged auto catalysts, suggesting that loss of activity is associated with sintering of the noble metals. Typical three-way catalysts used with engines operated with a stoichiometric air/fuel ratio are prepared by impregnating supports with noble metal solutions and heating to temperatures of about 450 to 700°C. This process does and is intended to decompose the noble metal compounds and leave the metals as separate crystallites on the support. In the present catalysts, which are operated under oxidizing conditions, it has been found that alloying the noble metals provides catalysts for reduction of nitrogen oxides.

10

15

20

25

30

Alloying may be done by various methods. In order to prepare supported catalysts in which Pt is alloyed with elements such as Co, Ni, Cu, Pd, Rh, Ag, or Au by high temperature thermal methods, the atmosphere must be sufficiently reducing that the alloyed elements are stable as the elements, rather than being stable in a higher oxidation state stabilized by the support lattice. Hydrogen may be used to carry out the reduction. When used, it will be present in more than zero up to 100 vol %, preferably about 1 to 97 vol % in the treating atmosphere. One preferred method of alloying is to thermally treat the impregnated support in an inert atmosphere containing up to 30 vol % steam, preferably about 10 vol % steam, and more than zero and up to 100 vol % hydrogen, at temperatures of about 600° to 1200° C selected to assure that any metal oxides are unstable with respect to the metals. Where the support is also produced by a process which uses a high temperature treatment to convert the alumina into a stable delta alumina, such as BaO-δ-Al2aO3 as discussed above, it is feasible to use such a process to prepare alloys of platinum. Thermal treatment with hydrogen present is needed particularly when the metals to be alloyed with Pt are Ni, Co, and Cu. For the other metals named, hydrogen is considered optional.

Hydrogen is not required when the metals are Pd, Rh, Ag, and Au. Alloys of Pt with Rh, Pd, Ag or Au may be prepared by exposing the metals to temperatures of  $600^{\circ}$  to  $1200^{\circ}$  C in an inert atmosphere, optionally containing 0-30 vol %  $H_2O$ .

Also, when hydrogen is absent oxygen may be introduced. The temperature and oxygen content of the gases are selected to provide the desired alloying. Alloys of Pt with Rh, Pd, and Au may be made in the presence of oxygen. In a specific example, Pt may be alloyed with Rh in an inert atmosphere which contains oxygen at temperatures below 1200° C at which Rh<sub>2</sub>O<sub>3</sub> is unstable with respect to Rh metal.

Generally, high temperature exposure to oxygen will increase the NO<sub>X</sub> reduction temperature window, as disclosed in co-pending USSN 08/130,340. Thus, the time and concentration of oxygen should be selected with this effect in mind since the alloying of Pt and the other metals is desired, but the peak temperature at which NO<sub>X</sub> reduction occurs will also be affected.

15

20

25

30

35

8

Supported alloy catalysts may also be prepared by colloidal methods. In this case, thermal treatment is not required. This method involves preparing a solution of a platinum compound, salt, or complex, together with a reducible compound, such as a salt, or a complex of the metal with which one desires to alloy the platinum. To this solution a reducing agent is added to reduce the metal ions to the zero valent state. Preferred reducing agents include sodium borohydride, hydrazine, citric acid, formic acid, sodium formate and amine-borane complex. Preferably, the rate of reduction of the platinum compound and the compound of the metal with which one desires to alloy with the platinum should proceed at roughly similar rates. Generally, a protecting agent such as a polycarboxylic acid, or a polymer, or a surfactant, or a multiply-charged ion, is added. Preferred protecting agents include citric acid, polyvinyl pyrolidone, polyethylene glycol, and compounds capable of producing a sulfite anion in water. The role of this protecting agent is to adsorb onto small metal particles as they form, hindering their growth. By this method, stable or metastable colloidal alloys of platinum may be dispersed in solution. After formation of the alloy colloid, the support material may be added as an additional sol or in powdered or granular form. Alternatively, the alloy colloid may be permitted to contact a catalyst support which has previously been applied to an inert monolithic carrier. Adsorption of the alloy onto the support material, followed by recovery of the support, drying, and optional calcination, completes the catalyst preparation.

The noble metals (which are also transition metals) of Groups 8-10 of the IUPAC Periodic Table consist of Pt, Pd, Rh, Ru, Os, and Ir. Generally, only Pt, Pd, and Rh are used in automobile exhaust catalysts. Ru, Os, and Ir easily form volatile oxides and, for this reason, are generally considered to be impractical, even for use in stoichiometric three-way catalysts. Stoichiometric three-way catalysts operate under conditions that are considerably more reducing than conditions encountered under lean-burn or diesel conditions. Although Ru, Os, and Ir may be alloyed with platinum by methods outlined herein, these alloys are not considered to be practically useful for purification of lean-burn or diesel exhaust gases. In addition to Pd and Rh, the transition metals Co, Ni, Cu, Ag, and Au can form alloys with platinum that modify the performance of platinum for NO<sub>x</sub> reduction under lean-burn or diesel conditions. The atomic ratio of Pt to Rh and Pd may range from 1/2 to 1000/1, preferably between 5/1 to 1000/1, most preferably from 5/1 to 50/1. The ratio of Pt to Co, Ni, and Cu may range

WO 95/32790 PCT/US95/06817

9

from 1/1 to 1000/1, preferably between 2/1 and 10/1. The ratio of Pt to Au may range from 15/1 to 1000/1, preferably 20/1 to 1000/1. The ratio of Pt to Ag may range from 1/10 to 1000/1. If disposed on a monolithic carrier, the Pt content may range from 0.3 g/L to 10 g/L, preferably 0.7 to 10 g/L, most preferably from 1 to 7 g/L. If disposed on a granular or spherical support, the platinum content may range from 0.15 wt.% to 7 wt.%, preferably between 0.33 wt.% and 5 wt.%, most preferably between 0.7 wt.% and 4 wt.%.

Optionally, base metals may be included in the catalysts as stabilizers or promoters provided they do not adversely affect the desired alloying of platinum and the other metals. If present, the base metals may range from about 0.01 to 50 g/L, most preferably 1 to 25 g/L based on a monolithic carrier. Addition of the base metals may be done by impregnation techniques familiar to those skilled in the art. Generally, these involve contacting the support with an aqueous solution of the metal (or metals) compound to be deposited and subsequently drying and heating the impregnated support to decompose the metal compounds.

The concentration of the metal compounds in solution will be adjusted to provide the amount of metals in the finished catalyst. The noble metal compounds may be those used in the past, such as halogenated compounds (e.g. chloro platinic acid, tetraamine platinum(II) chloride, rhodium chloride), nitrates (e.g. tetraamine platinum(II) nitrate, rhodium nitrate), acetates (e.g. rhodium acetate), sulfites, such as hydroxy disulfite platinum II acid, rhodium sulfite acid, and oxalates such as tetramethyl ammonium salts of bis-oxalato platinum(II) and tris-oxalato rhodium(III). Preferred noble metal compounds include rhodium nitrate, rhodium sulfite, rhodium oxalate, platinum oxalate and platinum sulfite.

The transition metals Co, Ni, Cu, Ag, and Au are preferably applied by impregnation of the support with an aqueous solution of soluble transition metal compounds. For gold, auric chloride or chloroauric acid may be used. For silver, silver nitrate, silver lactate, silver sulfite, or silver fluoride may be used. Compounds of cobalt, nickel, and copper may be chosen from the nitrate, acetate, lactate, citrate, or chloride. The choice of transition metal compounds will be determined by their kinetic ability to alloy with platinum.

30

10

15

20

25

PCT/US95/06817

10

15

20

25

30

35

#### Example 1

250 mL of a boehmite sol (VISTA Chemical Dispal 11N7-12) was placed in a 4000 mL beaker. 15.82 g barium acetate was dissolved in 10.3 mL of deionized water. The boehmite sol was stirred and the barium acetate solution along with 40.5 g of polyethylene glycol (ave. m.w. 400, Sigma PEG3265) were added simultaneously, producing a gel. The gel was transferred to an evaporating dish and dried in an oven at 120°C. The dried gel was then calcined at 1000°C for 6 hours. A pure δ alumina stabilized by barium, which we name BaO-δ-Al<sub>2</sub>O<sub>3</sub>, was the result, with no evidence of separate barium-containing phases.

9.45 g of platinum sulfite acid solution (2.55 wt.% Pt), 2.44 g of rhodium nitrate solution (0.50 wt.% Rh), and 0.036 g of cobalt (II) nitrate hexahydrate were dissolved in 170 g of deionized water. The solution was brought into contact with 18 g of the BaO-δ-Al<sub>2</sub>O<sub>3</sub> support prepared above. The resulting impregnated support was dried at 100°C for 3 hours and then calcined at 600°C for 6 hours. The calcined catalyst was hydrothermally treated at a temperature of about 1140°C for 6 hours in an atmosphere of nitrogen containing about 10% steam to alloy the noble metals. This Pt, Rh, Co/BaO-δ-Al<sub>2</sub>O<sub>3</sub> catalyst contained 1.32 wt.% Pt, 0.067 wt.% Rh, and 0.04 wt.% Co. A similar Pt/BaO-δ-Al<sub>2</sub>O<sub>3</sub> catalyst was prepared in the same manner except that the Rh and Co were omitted.

#### Example 2

A. 15.8 g of platinum sulfite acid solution (1.71 wt.% Pt) and 2.68 g of rhodium nitrate solution (0.50 wt.% Rh) were dissolved in 45 g of deionized water. The solution was brought into contact with 20 g of α-alumina made by calcining at 1350°C for 9 hours a commercial γ-alumina supplied by UOP. The resulting impregnated support was dried at 100°C for 3 hours. The dried catalyst was hydrothermally treated at a temperature of about 1194°C for 6 hours in an atmosphere of nitrogen containing about 10% steam to alloy the noble metals. This Pt,Rh/α-Al<sub>2</sub>O<sub>3</sub> catalyst contains 1.32 wt.% Pt and 0.067 wt.% Rh.

B. The preparation described above (A) was repeated except that 0.034 g of cobalt acetate tetrahydrate was dissolved in the platinum-rhodium solution so that the resulting catalyst contained WO 95/32790 PCT/US95/06817

11

cobalt in addition to the precious metals. This Pt,Rh, Co/α-Al<sub>2</sub>O<sub>3</sub> catalyst contained 1.36 wt.% Pt, 0.067 wt.% Rh, and 0.040 wt.% Co.

#### 5 Example 3

10

15

20

25

30

35

The four catalysts of Examples 1, 2A, and 2B were tested by exposure to a synthetic exhaust gas modeling emissions from a gasoline fueled lean-burn engine operating with an air-fuel ratio of about 18 which contained 1200 ppmv NO<sub>x</sub>, 3000 ppmv CO, 1000 ppmv H<sub>2</sub>, 1600 ppmv propene, 20 ppmv SO<sub>2</sub>, 10 vol.% steam, 10 wt.% CO<sub>2</sub>, and 3.2 vol.% O<sub>2</sub>, remainder nitrogen. The chemical equivalence ratio, expressed as the ratio of oxidizing chemical equivalents to reducing chemical equivalents is 3.54. The temperature was gradually increased up to a maximum of 800°C (rise 1) and then reduced to 50°C and repeated (rise 2). The performance of the catalysts is shown in Figure 1 for rise 1. It can be seen that each of the four catalysts exhibits an onset temperature for NO<sub>x</sub> reduction, a temperature of maximum NO<sub>x</sub> reduction, and a temperature above which NO<sub>x</sub> reduction does not occur. This contrasts with the performance of these same catalysts for oxidation of CO and hydrocarbons. Once a sufficient minimum temperature is attained, CO and hydrocarbon conversions increase with temperature to high, nearly quantitative, levels which do not decline substantially as temperature is increased to 800°C. Pt,Rh,Co/BaO-δ-Al<sub>2</sub>O<sub>3</sub> catalyst is considerably more effective than the Pt/BaOδ-Al<sub>2</sub>O<sub>3</sub> catalyst; this improvement is attributed to an alloyed nature of the Pt,Rh,Co catalyst. The modest improvement of the Pt,Rh,Co/α-Al<sub>2</sub>O<sub>3</sub> compared to the Pt,Rh/\alpha-Al2O3 is attributed to incorporation of Co into the Pt,Rh alloyed crystallites.

#### Example 4

A catalyst was prepared by applying a solution of chloroplatinic acid and rhodium chloride to 20-40 mesh pellets of a CeO<sub>2</sub>-Al<sub>2</sub>O<sub>3</sub> (24 wt.% CeO<sub>2</sub>), followed by evaporation and calcination in air at 600°C. Analysis revealed the Pt content to be 4340 ppm by weight and the Rh content to be 1340 ppm by weight. In this form, the catalyst is not alloyed. The Pt and Rh in the catalyst were alloyed by treatment at 1200°C in flowing N<sub>2</sub> with 10% steam. The unalloyed and alloyed catalysts were tested using the gas composition and temperature program used in Example 3. Figure 2 shows performance of these

two catalysts both during rise 1 and rise 2. The alloyed catalyst exhibits superior NO reduction performance.

Figure 3 shows the 311 reflection of the Pt,Rh catalyst as examined by X-ray diffraction using Cu-Kα radiation. If substantial amounts of pure Pt crystallites were present, an intensity feature centered at 81.28° 2θ would occur; similarly if substantial amounts of pure Rh crystallites were present, an intensity feature centered at 84.39° 2θ would occur. Unfortunately, α-Al<sub>2</sub>O<sub>3</sub> also has a reflection centered at 84.37° 2θ, rendering detection of any pure Rh impossible. Substantial diffraction intensity due to Pt,Rh alloys is apparent between about 81.3° 2θ and 83.0° 2θ. A range of Pt,Rh compositions appears to occur, rendering any attempts to determine crystallite size by line shape analysis meaningless and, indeed, erroneous. Figures 2 and 3 demonstrate Pt,Rh alloys are capable of reducing NO<sub>X</sub> under highly oxidizing, lean-burn conditions.

15

20

25

30

10

5

#### Example 5

A series of catalysts was prepared using a delta alumina manufactured by Condea, an aqueous solution of chloroplatinic acid, and an aqueous solution of cupric acetate monohydrate. After evaporation of the solution onto the alumina support, the material was calcined in air at 600°C. The Pt content was 1.32 wt.%, the Cu content 2000 ppm by weight. Portions of this catalyst were treated for six hours at the following conditions: 10% steam and 90% N<sub>2</sub> at 1140°C (Treatment A), 10% steam and 90% N<sub>2</sub> at 1200°C (Treatment B), and 10% steam, 85% N<sub>2</sub>, and 5% H<sub>2</sub> at 1200°C (Treatment C). The catalysts made by Treatments A, B, and C were examined by X-ray diffraction to determine the extent of alloying between Pt and Cu. The X-ray diffraction reflections for the metal occurred at angles greater than those for corresponding Pt-only reflections but lower than those expected for a stoichiometric alloy where all the copper is alloyed with all the platinum. The extent of alloying varied among the three treatments. We determined the apparent unit cell sizes for the three Pt,Cu catalysts; from this the mole fraction of Copper present in the Pt,Cu alloys can be determined. The following are the results of this analysis:

15

20

25

30

	Unit cell size	Mole fraction Cu
Treatment A	3.854 Å	22.4%
Treatment B	3.850 Å	23.6%
Treatment C	3.844 Å	25.7%

Separate reflections due to pure Pt or pure Cu are not observed. It is believed that all the Pt is metallic while only a fraction of Cu occurs in metallic form alloyed with the Pt while the rest of the Cu is dispersed on the alumina as oxidized cupric species. The more reducing the treatment, the greater the fraction of copper present in the alloys produced.

In addition, an additional Pt,Cu catalyst containing only 700 ppm Cu was prepared using the same alumina provided by Condea, the same preparative method, and subjected to Treatment C. It was not subjected to X-ray diffraction analysis, but by analogy we expect a fraction of the copper is alloyed with the platinum.

For comparison, three Pt-only catalysts were prepared using the same alumina provided by Condea, the same preparative method, and subjected to Treatment B, Treatment C, or a Treatment D which involved 10% H<sub>2</sub>0, 90% N<sub>2</sub>. at 1100°C.

These catalysts were evaluated using a synthetic gas blend that models exhaust from a gasoline lean-burn engine operating at an Air-Fuel Ratio of 22. This gas contains 1000 ppmv CO, 330 ppmv H2, 3300 ppmv propene, 700 ppmv NO, 7.5 vol.% O2, 10 vol.% CO2, 10 vol.% H2O, and balance N2; the flow rate was 5 liters/minute. The chemical equivalence ratio of this gas mixture is 5.34. The reactor inlet temperature was increased from 150°C to 800°C at 5°C/minute. NO<sub>x</sub> conversion commences at temperatures near those where propene conversion commences. In all cases, once NO<sub>X</sub> conversion commences, it increases monotonically with increasing temperature to a temperature of maximum NOx conversion; above this temperature, NO<sub>X</sub> conversion decreases monotonically. Performance may be compared by noting the maximum NO conversion level, the temperature of maximum NO<sub>X</sub> conversion, and by comparing relative total amounts of NO reduced. The total amount of NO reduced may be determined by integrating the observed time dependence of NO<sub>X</sub> conversion with respect to time. We define the integral amount of NO reduced by the Pt,Cu catalysts containing 2000 ppm Cu and subjected to Treatment C as 100. Performance of the Pt,Cu and Pt-only catalysts is summarized immediately below.

Amount of Cu	<u>Treatment</u>	Max. NO	Temp. of Max.	Integral NO
		Conv.	NO Conv.	
2000 ppm	С	48%	219°C	100
700 ppm	С	40%	238°C	99
2000 ppm	В	13%	224 °C	47
2000 ppm	A	18%	226°C	54
0	С	24%	297 °C	55
0	В	24%	280 °C	63
0	D	26%	268 °C	75

The results of the foregoing table demonstrate the Pt,Cu catalysts to possess properties different from those of the Pt-only catalysts. The Pt,Cu catalysts prepared with reduction at 1200°C reduce the most NO of the Pt,Cu catalysts while among the Pt-only catalysts, the catalyst subjected to this treatment reduces the least amount of NO. Larger amounts of Cu results in maximum NO conversion at lower temperatures. The most alloyed catalyst reduces the most NO. Alloyed Pt,Cu catalysts reduce NO at temperatures lower than those of similarly prepared Pt-only catalysts.

The Pt,Cu catalyst with 2000 ppm Cu that was subjected to treatment C together with the Pt-only catalysts subjected to treatment C were tested using the synthetic gas blend of this example with the addition of 20 ppm SO<sub>2</sub>. The results of these two tests are summarized below. The alloyed Pt,Cu catalyst reduces more NO than the Pt-only catalyst, and the temperature of maximum NO reduction is nearly 100°C lower than that of the Pt-only catalyst.

	Max. NO Conv.	Temp. of Max.	Integral NO
		NO Conv.	
Pt,Cu	31%	249°C	73
Pt-only	17%	344°C°	33

20

15

Alloying Pt with Cu substantially changes the nature of catalytic NO reduction under lean-burn conditions compared to Pt alone. Typically, they are effective at lower temperatures than similar Pt-only catalysts. Although

WO 95/32790

15

PCT/US95/06817

performance at lower temperatures is generally considered to be a benefit in catalysis, this is not necessarily the case for lean NO<sub>X</sub> catalysts. Since lean NO<sub>X</sub> catalysts exhibit NO<sub>X</sub> control over characteristic ranges of temperatures, and do not reduce NO<sub>X</sub> at lower or higher temperatures outside of their characteristic range, lower temperature performance is not necessarily advantageous. Indeed, control of NO<sub>X</sub> emissions from lean-burn or diesel engines will be needed over a range of temperatures greater than the effective temperature range of any single catalyst considered in this invention. As such, sets of appropriately situated catalysts each possessing different temperature ranges over which they are effective will be required in order to provide adequate control of NO<sub>X</sub> emissions. The Integral NO values for these catalysts shows Pt catalysts alloyed with Cu can be more effective at reducing NO in the temperature window where they are effective under lean-burn conditions than Pt-only catalysts prepared in the same manner are in the temperature window where Pt-only are effective.

15

20

25

30

10

#### Example 6

A similar series of Pt, Ag catalysts containing 1.32% Pt, 2000 ppm Ag catalysts was prepared using the same alumina support manufactured by Condea as used in Example 5 and a solution of chloroplatinic acid and silver nitrate. The solution was evaporated to deposit the metal complexes onto the alumina, the resulting material calcined at 600°C, and portions of the calcined material subjected to treatments A, B, and C to prepare three catalysts. The catalysts were examined by X-ray diffraction. Reflections due to Pt-only crystallites and due to Pt, Ag alloyed crystallites show that in contrast to the Pt, Cu catalysts, the alloyed crystallites in the three Pt, Ag catalysts have the same composition which is Ag-rich, rather than Pt-rich, containing, on average, 87% Ag. The intensity of the alloyed Pt, Ag reflections are considerably greater in the catalyst made with treatment C compared to those made by treatments A and B. This suggests that either the Pt,Ag crystallites are more highly sintered in the catalyst made by treatment C or that the mass fraction of Pt and Ag present in alloyed crystallites is greatest in the catalyst made by treatment C. The catalysts were evaluated by the method of Example 5 using the gas mixture that did not contain SO2; results are summarized in the table immediately below.

10

15

20

25

30

Treatment	Max. NO Conv.	Temp. of Max.	Integral NO
		NO Conv.	
A	20%	31 <b>8 °C</b>	47
В	23%	272 °C	46
С	43%	261℃	95

The catalyst made with treatment C was also tested by the method of Example 5 using a gas mixture that contained 20 ppmv SO<sub>2</sub>. The maximum NO conversion was 21%; the temperature of maximum NO conversion 270°C; the integral NO value 70. The Pt,Ag catalyst made by treatment C reduces considerably more NO than either the other Pt,Ag catalysts or the Pt-only catalysts of Example 5. We attribute the superior performance of the Pt,Ag catalyst made with treatment C to the probability that the alloyed crystallites dominate catalytic performance. The temperature range in which the Pt,Ag catalysts are effective varies considerably more than analogous Pt-only catalysts. The more forcing the alloying condition, the lower the range of effective temperatures for NO reduction. In contrast, the more forcing the treatment condition for the Pt-only catalysts, the higher the range of effective temperatures for NO reduction. As was the case for the Pt,Cu catalysts, alloying Pt with Ag modifies considerably the performance of the resulting catalyst for NO reduction under lean-burn conditions.

#### Example 7

A series of Pt,Rh alloyed catalysts was prepared supported on alpha alumina. 55.57 grams Pt sulfite solution containing 2.55 wt.% Pt and 54.33 g Rh nitrate solution containing 0.50 wt.% Rh were diluted with 307.75 grams deionized water. 140 grams α-Al<sub>2</sub>O<sub>3</sub> was added and the solution was evaporated onto the alumina over a course of three hours. The catalyst contains 1.0 wt.% Pt and 0.19 wt.% Rh. Portions of these catalysts were treated at about 1050°C in atmospheres containing 10 vol.% steam and various amounts of oxygen; i.e., 0 vol.% O<sub>2</sub>, 200 ppm O<sub>2</sub>, 600 ppm O<sub>2</sub>, 1800 ppm O<sub>2</sub>, 4800 ppm O<sub>2</sub>, or 21 vol.% O<sub>2</sub>; the balance was N<sub>2</sub>. These materials were examined by X-ray diffraction. The material treated with 10% steam, 0% O<sub>2</sub>, and 90% N<sub>2</sub> contained a broad, apparently bimodal, distribution of alloyed crystallites, with maxima of diffraction intensity corresponding to crystallites that are relatively Pt-

15

20

30

rich (3.9 atom% Rh) and relatively Rh-rich (30.7 atom% Rh). A stoichiometric alloy wherein all the Rh is alloyed with all the Pt would contain 26.3 atom% Rh. The catalysts treated with O<sub>2</sub> partial pressures of 200 ppm or greater all contained unimodal distributions of Pt,Rh alloys. The average Rh content, as determined by X-ray diffraction analysis, of the alloyed crystallites in these catalysts is summarized below.

200 ppm O <sub>2</sub>	22.7 atom% Rh
600 ppm O <sub>2</sub>	23.2 atom% Rh
1800 ppm O <sub>2</sub>	23.6 atom% Rh
4800 ppm O <sub>2</sub>	24.5 atom% Rh
21% O <sub>2</sub>	20.6 atom% Rh

For low partial pressures of  $O_2$ , the extent of alloying increases with  $O_2$  partial pressure, but the extent of alloying is lower when air, with 21%  $O_2$ , was used. The important point is that alloys were formed in all these samples. This is due, in part, to the fact that  $Rh_2O_3$  decomposes in air at temperatures of about  $1033^{\circ}C$ , so Rh metal is favored at treatment temperatures. Transport of Pt and Rh via unstable, volatile oxides, may facilitate alloy formation.

These five catalysts were tested using the gas mixture and temperature program of Example 3. Each catalyst exhibited a temperature window or range of temperatures where NO was reduced. The maximum NO conversion was between 35-40% for each of these catalysts. The temperature of maximum NO conversion occurred between 200°C and 325°C; the greater the O<sub>2</sub>, partial pressure during the treatment that formed the alloys, the higher the temperature of maximum NO conversion. At the 20% conversion level, the temperature windows for NO conversion varied in width between about 50°C to about 70°C.

#### 25 Example 8

Alloyed catalysts of the present invention may be made by application of colloidal alloys of platinum onto suitable support materials or onto sol precursors of suitable support materials. A series of Pt,Rh/SiO<sub>2</sub> materials were prepared since silica (specifically, Cab-O-Sil M5) is a structurally amorphous material that provides a highly favorable substrate for characterization by X-ray diffraction. Materials containing about 2.7 wt% Pt

15

20

25

were prepared with a theoretical atomic% Pt relative to the sum of Pt and Rh varying between 57% and 90%. Solutions containing suitable amounts of chloroplatinic acid and rhodium chloride were prepared to which citric acid monohydrate was added. The mole ratio of citric acid to platinum was about 10; the role of the citric acid is to stabilize the Pt,Rh alloy colloid particles with respect to growth and precipitation. Then a stoichiometric excess of a sodium borohydride solution was rapidly added. Rapid reaction ensued, and the solution turned black, consistent with production of a colloidal metal. The silica was added and the solution evaporated to dryness. The resulting materials were examined by X-ray diffraction; the apparent Pt unit cell lattice constant determined, and the amount of Pt present in the crystallites determined. Results are summarized in the Table immediately below:

Alloy Lattice	Theoretical Pt	Experimental Pt
Parameter (Å)	atomic %	atomic %
3.905	90	85
3.891	83	74
3,880	77	64
3.881	66	65
3.855	57	52

The results show that the colloidal material formed by the sodium borohydride reduction consists of alloyed particles of Pt and Rh. The reflections were broad, suggesting that a range of compositions may have occurred.

A similar preparation provides a material that is catalytically active for the desired reaction. A reducing solution was made by dissolving 5.6 g NaOH in 400 g deionized H<sub>2</sub>O; 4.0 g NaBH<sub>4</sub> was added to this solution. A precious metal solution containing 0.28 mg Rh and 26 mg Pt and about 903 g deionized water was prepared in a 4 liter beaker, 40 g polyvinylpyrollidone (PVP) solution (1 g PVP/1 water) was added to this precious metal solution. The PVP serves to stabilize the metal colloids after they form, enhancing their stability with respect to growth and precipitation. The reducing solution was rapidly added to the precious metal solution with stirring; bubbles due to H<sub>2</sub> formation formed rapidly and provided evidence that the reduction proceeded. A second solution Pt,Rh colloid was prepared in an identical manner. These two batches of Pt,Rh colloid were allowed to sit overnight. The batches of Pt,Rh

WO 95/32790 PCT/US95/06817

19

colloid were combined in a 4-liter beaker and 115.4 g of Vista DISPAL 11N7-12 alumina sol was added. The precious metal adsorbed onto the alumina sol, which precipitated. After settling, the supernatant was decanted, the resulting solid recovered and dried at 500°C for 6 hr in flowing N<sub>2</sub>. The DISPAL solution contains colloidal boehmite, and the drying temperature was chosen to be sufficiently high to cause the boehmite to transform to γ-Al<sub>2</sub>O<sub>3</sub>. The alloyed catalyst contained about 0.7 wt.% Pt and about 0.035 wt.% Rh.

This catalyst was then tested using the gas mixture of Example 3. The Pt,Rh alloy colloid catalyst of the present example may be compared with the Pt,Rh,Co/BaO-δ-Al<sub>2</sub>O<sub>3</sub> alloyed catalyst of Example 1. The Pt,Rh alloy colloid exhibits an onset of NO reduction at about 250-275°C while the Pt,Rh,Co/BaO-δ-Al<sub>2</sub>O<sub>3</sub> exhibits an onset of NO reduction at about 325-350°C. The two catalysts reduce substantially the same amount of NO over the temperature rise (to within about 1.3%) although the Pt,Rh,Co/BaO-δ-Al<sub>2</sub>O<sub>3</sub> reduces NO over a temperature range of about 150°C while the Pt,Rh colloid catalyst reduces NO over a temperature range of about 250-275°C.

10

15

10

15

35

#### **CLAIMS**:

- A catalyst for removing carbon monoxide, hydrocarbons, and nitrogen oxides from the exhaust of an internal combustion engine containing oxygen in substantial excess of the stoichiometric quantity needed for complete combustion consisting essentially of platinum alloyed with at least one metal selected from the group consisting of cobalt, nickel, copper, rhodium, palladium, silver, and gold on an inert support.
- A catalyst of Claim 1 wherein said inert support is a temperature stable support selected from the group consisting of barium-stabilized alumina, α-alumina, δ-alumina, θ-alumina, zirconia, titania, silicon carbide, and cordierite.
- 3. A catalyst of Claim 2 wherein platinum is alloyed with at least one member of the group consisting of cobalt, nickel, copper, palladium, rhodium, silver and gold by heating in an inert atmosphere containing 0-30 vol. % steam, greater than zero up to 100 vol.% hydrogen, and zero up to less than 100 vol.% inert gas at temperatures in the range of about 600° to 1200°C.
- 4. A catalyst of Claim 2 wherein platinum is alloyed with at least one member of the group consisting of palladium, rhodium, silver, and gold by heating in an inert gas containing up to 30 vol.% steam at temperatures in the range of about 600° to 1200°C.
  - 5. A catalyst of Claim 2 wherein said temperature stable support is  $\alpha$ -Al<sub>2</sub>O<sub>3</sub>.
- 6. A catalyst of Claim 5 wherein platinum is alloyed with rhodium by heating in an atmosphere containing 0-30 vol.% steam, 0-21 vol.% oxygen and an inert gas at a temperature below 1200°C at which Rh<sub>2</sub>O<sub>3</sub> is unstable with respect to Rh metal.
- 7. A catalyst of Claim 2 wherein platinum is alloyed with at least one member of the group consisting of palladium, rhodium, and gold by heating in an atmosphere containing up to 21 vol.% oxygen, 0-30 vol.% stream and an inert gas at temperatures in the range of about 600° to 1200°C.
  - 8. A process for preparing a catalyst containing platinum alloyed with at least one metal selected from the group consisting of cobalt, nickel, copper, rhodium, palladium, silver, and gold comprising

15

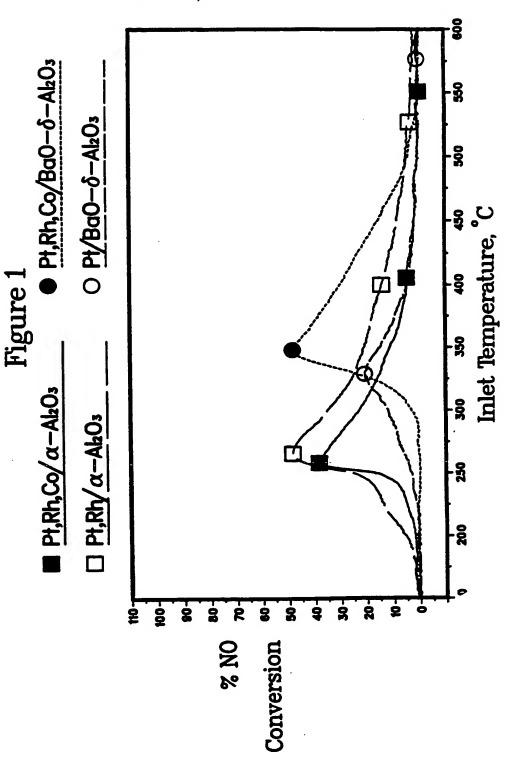
WO 95/32790 PCT/US95/06817

(a) reacting in a solvent soluble platinum compounds and a soluble compound of at least one of said metal in the presence of a reducing agent capable of producing a colloidal suspension of platinum alloyed with said metals;

21

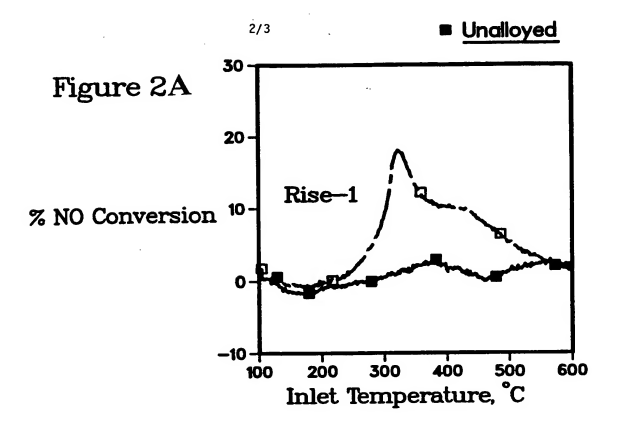
- (b) adsorbing said colloidal suspension produced in (a) onto a support;
  - (c) drying and optionally calcining said support of (b);
  - (d) recovering the dried support of (c).
- 9. The process of Claim 8 wherein said reducing agent is selected from the group consisting of sodium borohydride, hydrazine, citric acid, formic acid, sodium formate, and amine-borane complex.
  - 10. The process of Claim 8 wherein said platinum compounds are selected from the group consisting of chloroplatinic acid, platinum(II) hydroxy sulfite acid, tetraamine platinum(II) chloride, and tetraamine platinum(II) nitrate.
  - 11. The process of Claim 8 wherein the reaction of step (a) is carried out in the presence of a protecting agent selected from the group consisting of citric acid, polyvinyl pyrolidone, polyethylene glycol, and a compound producing a sulfite anion in water.
- The process of Claim 8 wherein said support is selected from the group consisting of (a) temperature stable supports consisting of barium-stabilized alumina, α-alumina, δ-alumina, θ-alumina, zirconia, titania, silicon carbide, and cordierite and (b) less temperature stable supports consisting of γ-alumina, crystalline boehmite, crystalline alumina silicates, non-crystalline silica alumina, α-Zr(HPO<sub>4</sub>)<sub>2</sub>, NaZr<sub>2</sub>(PO<sub>4</sub>)<sub>3</sub>, and (Na,H)Zr<sub>2</sub>(PO<sub>4</sub>)<sub>3</sub>.

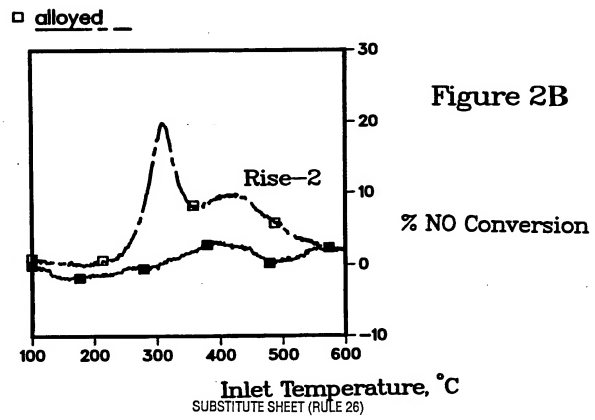


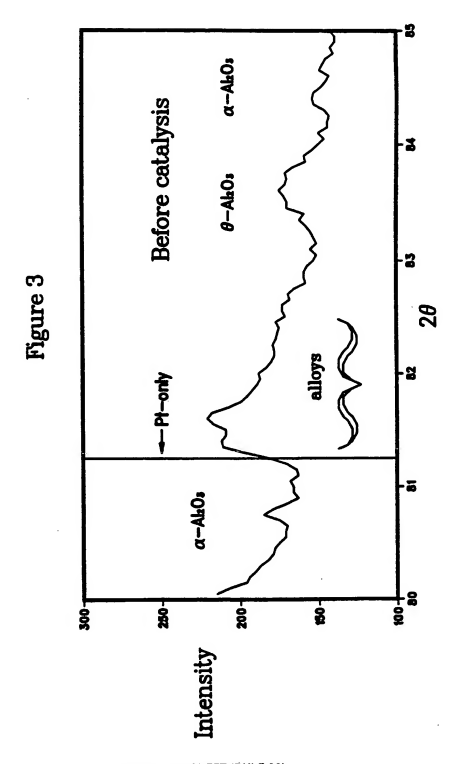


SUBSTITUTE SHEET (RULE 26)

WO 95/32790 PCT/US95/06817







SUBSTITUTE SHEET (RULE 26)

# INTERNATIONAL SEARCH REPORT Inter tal Application No

Inter 1al Application No PCT/US 95/06817

A. CLASS IPC 6	B01D53/94 B01J23/38 B01J23/	89	
According	to International Patent Classification (IPC) or to both national classi	fication and IPC	
B. FIELD	S SEARCHED		
Minimum of IPC 6	documentation searched (classification system followed by classification by BO1D BO1J	tion symbols)	
	tion searched other than minimum documentation to the extent that		
Electronic o	data base consulted during the international search (name of data ba	se and, where practical, search terms used)	
C. DOCUM	MENTS CONSIDERED TO BE RELEVANT		
Category *	Citation of document, with indication, where appropriate, of the r	elevant passages	Relevant to claim No.
P,X	WO,A,95 09687 (ALLIED SIGNAL INC. April 1995		1-7
	see page 7, line 19 - page 9, line claims 1-6; examples	ne 29;	
Y			8-12
х	FR,A,2 130 310 (JOHNSON MATTHEY & LTD.) 3 November 1972 see page 13, line 10 - page 14, claims 1-15; example		1-3,5
x	FR,A,2 321 940 (ENGELHARD MINERAL CHEMICALS CORP.) 25 March 1977 see claims 1-19; examples 1-5	LS &	1,2
		-/	
	-		
X Fur	ther documents are listed in the continuation of box C.	χ Patent family members are listed	in annex.
* Special ca	ategories of cited documents :	"T" later document published after the into or priority date and not in conflict we cited to understand the principle or t	ith the application but
	dered to be of particular relevance document but published on or after the international date	"X" document of particular relevance; the cannot be considered novel or canno	: claimed invention
"L" docum	nent which may throw doubts on priority daim(s) or is cited to establish the publication date of another on or other special reason (as specified)	involve an inventive step when the d "Y" document of particular relevance; the cannot be considered to involve an in	ocument is taken alone claimed invention
O' docum	nent referring to an oral disclosure, use, exhibition or means	document is combined with one or n ments, such combination being obvious in the art.	nore other such docu-
later t	nent published prior to the international filing date but than the priority date claimed	*&* document member of the same patern  Date of mailing of the international s	
	e actual completion of the international search		•
	21 September 1995	27.09.199!	· · · · · · · · · · · · · · · · · · ·
Name and	mailing address of the ISA  European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk	- American amang	
	Tel. (+31-70) 340-2040, Tx. 31 651 epo nl,	Eijkenboom, A	

# INTERNATIONAL SEARCH REPORT

Interr. 1al Application No PCT/US 95/06817

		PCT/US 95/06817		
C.(Continua	n) DOCUMENTS CONSIDERED TO BE RELEVANT			
Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.		
X	US,A,4 025 606 (JOHNSON & MATTHEY & CO. LTD.) 24 May 1977 see column 6, line 23 - column 14, line 21; claims 1-23	1-3,5		
Y	US,A,4 152 301 (GENERAL MOTORS CORP.) 1 May 1979 see column 7, line 20 - line 62; claims 1-12	8-12		
A	US,A,4 791 091 (ALLIED-SIGNAL INC.) 13 December 1988 see column 7, line 14 - line 66; claims 1-20	1,2,8,		

# INTERNATIONAL SEARCH REPORT

information on patent family members

Intern ul Application No
PCT/US 95/06817

WO-A-9509687 13-04-95 NONE  FR-A-2130310 03-11-72 GB-A- 1390182 09-04-75 BE-A- 780767 17-07-72 CA-A- 1002731 04-01-77 DE-C- 2212616 12-08-93 DE-A- 2212616 28-09-72 NL-A- 7203388 19-09-72 SE-B- 428761 25-07-83 US-A- 3840471 08-10-74					
FR-A-2130310 03-11-72 GB-A- 1390182 09-04-75 BE-A- 780767 17-07-72 CA-A- 1002731 04-01-77 DE-C- 2212616 12-08-93 DE-A- 2212616 28-09-72 NL-A- 7203388 19-09-72 SE-B- 428761 25-07-83 US-A- 3840471 08-10-74  FR-A-2321940 25-03-77 AU-B- 505606 29-11-79 AU-B- 1713376 02-03-78 BE-A- 845625 28-02-77 CA-A- 1083124 05-08-80 DE-A- 2637198 03-03-77 GB-A- 1561683 27-02-80 JP-A- 52027088 01-03-77 SE-B- 446506 22-09-86 SE-A- 7609423 28-02-77 US-A- 4552733 12-11-85 US-A- 4157316 05-06-79  US-A-4025606 24-05-77 GB-A- 1405405 10-09-75 DE-A- 2231296 19-04-73 US-A- 3839225 01-10-74  US-A-4152301 01-05-79 AU-B- 524724 30-09-82 AU-B- 3803978 17-01-80 CA-A- 1098105 24-03-81 DE-A- 2830686 01-02-79 GB-A, B 2000985 24-01-79 JP-A- 54019494 14-02-79 US-A-4791091 13-12-88 EP-A, B 0310398 05-04-89 JP-T- 3501355 28-03-91 WO-A- 8902782 06-04-89	Patent document cited in search report				
BE-A	WO-A-9509687	13-04-95	NONE		
CA-A- 1002731 04-01-77 DE-C- 2212616 12-08-93 DE-A- 2212616 28-09-72 NL-A- 7203388 19-09-72 SE-B- 428761 25-07-83 US-A- 3840471 08-10-74  FR-A-2321940 25-03-77 AU-B- 505606 29-11-79 AU-B- 1713376 02-03-78 BE-A- 845625 28-02-77 CA-A- 1083124 05-08-80 DE-A- 2637198 03-03-77 GB-A- 1561683 27-02-80 JP-A- 52027088 01-03-77 SE-B- 446506 22-09-86 SE-A- 7609423 28-02-77 US-A- 4157316 05-06-79 US-A- 4157316 05-06-79 US-A- 2231296 19-04-73 US-A- 2231296 19-04-73 US-A- 2331296 19-04-73 US-A- 1098105 24-03-81 DE-A- 2830686 01-02-79 GB-A, B 2000985 24-01-79 JP-A- 54019494 14-02-79 US-A-4791091 13-12-88 EP-A, B 0310398 05-04-89 JP-T- 3501355 28-03-91 WO-A- 8902782 06-04-89	FR-A-2130310	03-11-72			
DE-C- 2212616 12-08-93 DE-A- 2212616 28-09-72 NL-A- 7203388 19-09-72 SE-B- 428761 25-07-83 US-A- 3840471 08-10-74  FR-A-2321940 25-03-77 AU-B- 505606 29-11-79 AU-B- 1713376 02-03-78 BE-A- 845625 28-02-77 CA-A- 1083124 05-08-80 DE-A- 2637198 03-03-77 GB-A- 1561683 27-02-80 JP-A- 52027088 01-03-77 SE-B- 446506 22-09-86 SE-A- 7609423 28-02-77 US-A- 4552733 12-11-85 US-A- 4552733 12-11-85 US-A- 4552733 12-11-85 US-A- 4552733 12-11-85 US-A- 3839225 01-10-74  US-A-4152301 01-05-79 AU-B- 524724 30-09-82 AU-B- 3803978 17-01-80 CA-A- 1098105 24-03-81 DE-A- 2830686 01-02-79 GB-A, B 2000985 24-01-79 JP-A- 54019494 14-02-79  US-A-4791091 13-12-88 EP-A, B 0310398 05-04-89 JP-T- 3501355 28-03-91 WO-A- 8902782 06-04-89					
DE-A-   2212616   28-09-72   NL-A-   7203388   19-09-72   SE-B-   428761   25-07-83   US-A-   3840471   08-10-74					
NL-A- 7203388					
SE-B- 428761 25-07-83 US-A- 3840471 08-10-74  FR-A-2321940 25-03-77 AU-B- 505606 29-11-79 AU-B- 1713376 02-03-78 BE-A- 845625 28-02-77 CA-A- 1083124 05-08-80 DE-A- 2637198 03-03-77 GB-A- 1561683 27-02-80 JP-A- 52027088 01-03-77 SE-B- 446506 22-09-86 SE-A- 7609423 28-02-77 US-A- 4552733 12-11-85 US-A- 4157316 05-06-79  US-A-4025606 24-05-77 GB-A- 1405405 10-09-75 DE-A- 2231296 19-04-73 US-A- 3839225 01-10-74  US-A-4152301 01-05-79 AU-B- 524724 30-09-82 AU-B- 3803978 17-01-80 CA-A- 1098105 24-03-81 DE-A- 2830686 01-02-79 JP-A- 54019494 14-02-79  US-A-4791091 13-12-88 EP-A, B 0310398 05-04-89 JP-T- 3501355 28-03-91 WO-A- 8902782 06-04-89					
US-A- 3840471 08-10-74  FR-A-2321940 25-03-77 AU-B- 505606 29-11-79 AU-B- 1713376 02-03-78 BE-A- 845625 28-02-77 CA-A- 1083124 05-08-80 DE-A- 2637198 03-03-77 GB-A- 1561683 27-02-80 JP-A- 52027088 01-03-77 SE-B- 446506 22-09-86 SE-A- 7609423 28-02-77 US-A- 4552733 12-11-85 US-A- 4157316 05-06-79  US-A-4025606 24-05-77 GB-A- 1405405 10-09-75 DE-A- 2231296 19-04-73 US-A- 3839225 01-10-74  US-A-4152301 01-05-79 AU-B- 524724 30-09-82 AU-B- 3803978 17-01-80 CA-A- 1098105 24-03-81 DE-A- 2830686 01-02-79 GB-A, B 2000985 24-01-79 JP-A- 54019494 14-02-79  US-A-4791091 13-12-88 EP-A, B 0310398 05-04-89 JP-T- 3501355 28-03-91 WO-A- 8902782 06-04-89					
FR-A-2321940 25-03-77 AU-B- 505606 29-11-79 AU-B- 1713376 02-03-78 BE-A- 845625 28-02-77 CA-A- 1083124 05-08-80 DE-A- 2637198 03-03-77 GB-A- 1561683 27-02-80 JP-A- 52027088 01-03-77 SE-B- 446506 22-09-86 SE-A- 7609423 28-02-77 US-A- 4552733 12-11-85 US-A- 4157316 05-06-79  US-A-4025606 24-05-77 GB-A- 1405405 10-09-75 DE-A- 2231296 19-04-73 US-A- 3839225 01-10-74  US-A-4152301 01-05-79 AU-B- 524724 30-09-82 AU-B- 3803978 17-01-80 CA-A- 1098105 24-03-81 DE-A- 2830686 01-02-79 GB-A, B 2000985 24-01-79 JP-A- 54019494 14-02-79  US-A-4791091 13-12-88 EP-A, B 0310398 05-04-89 US-A-8902782 06-04-89					
AÜ-B- 1713376 02-03-78 BE-A- 845625 28-02-77 CA-A- 1083124 05-08-80 DE-A- 2637198 03-03-77 GB-A- 1561683 27-02-80 JP-A- 52027088 01-03-77 SE-B- 446506 22-09-86 SE-A- 7609423 28-02-77 US-A- 4552733 12-11-85 US-A- 4552733 12-11-85 US-A- 4157316 05-06-79  US-A- 4025606 24-05-77 GB-A- 1405405 10-09-75 DE-A- 2231296 19-04-73 US-A- 3839225 01-10-74  US-A-4152301 01-05-79 AU-B- 524724 30-09-82 AU-B- 3803978 17-01-80 CA-A- 1098105 24-03-81 DE-A- 2830686 01-02-79 GB-A, B 2000985 24-01-79 JP-A- 54019494 14-02-79  US-A-4791091 13-12-88 EP-A, B 0310398 05-04-89 US-A- 8902782 06-04-89					
BE-A-	FR-A-2321940	25-03-77			
US-A-4025606 24-05-77 GB-A- 1405405 10-09-75 US-A-4152301 01-05-79 AU-B- 3803978 17-01-80 CA-A- 1098105 24-03-81 DE-A- 2830686 01-02-79 GB-A, B 2000985 24-01-79 JP-A- 54019494 14-02-79 US-A-4791091 13-12-88 EP-A, B 0310398 05-04-89 US-A- 8902782 06-04-89					
DE-A- 2637198 03-03-77 GB-A- 1561683 27-02-80 JP-A- 52027088 01-03-77 SE-B- 446506 22-09-86 SE-A- 7609423 28-02-77 US-A- 4552733 12-11-85 US-A- 4157316 05-06-79  US-A-4025606 24-05-77 GB-A- 1405405 10-09-75 DE-A- 2231296 19-04-73 US-A- 3839225 01-10-74  US-A-4152301 01-05-79 AU-B- 524724 30-09-82 AU-B- 3803978 17-01-80 CA-A- 1098105 24-03-81 DE-A- 2830686 01-02-79 GB-A, B 2000985 24-01-79 JP-A- 54019494 14-02-79  US-A-4791091 13-12-88 EP-A, B 0310398 05-04-89 JP-T- 3501355 28-03-91 WO-A- 8902782 06-04-89					
GB-A- 1561683 27-02-80 JP-A- 52027088 01-03-77 SE-B- 446506 22-09-86 SE-A- 7609423 28-02-77 US-A- 4552733 12-11-85 US-A- 4157316 05-06-79  US-A- 4157316 05-06-79  US-A- 2231296 19-04-73 US-A- 3839225 01-10-74  US-A-4152301 01-05-79 AU-B- 524724 30-09-82 AU-B- 3803978 17-01-80 CA-A- 1098105 24-03-81 DE-A- 2830686 01-02-79 GB-A, B 2000985 24-01-79 JP-A- 54019494 14-02-79  US-A-4791091 13-12-88 EP-A, B 0310398 05-04-89 JP-T- 3501355 28-03-91 WO-A- 8902782 06-04-89					
US-A-4025606 24-05-77 GB-A- 1405405 10-09-75 US-A- 4157316 05-06-79  US-A-4025606 24-05-77 GB-A- 1405405 10-09-75 DE-A- 2231296 19-04-73 US-A- 3839225 01-10-74  US-A-4152301 01-05-79 AU-B- 3803978 17-01-80 CA-A- 1098105 24-03-81 DE-A- 2830686 01-02-79 GB-A, B 2000985 24-01-79 JP-A- 54019494 14-02-79  US-A-4791091 13-12-88 EP-A, B 0310398 05-04-89 JP-T- 3501355 28-03-91 WO-A- 8902782 06-04-89					
SE-B- 446506 22-09-86 SE-A- 7609423 28-02-77 US-A- 4552733 12-11-85 US-A- 4157316 05-06-79  US-A-4025606 24-05-77 GB-A- 1405405 10-09-75 DE-A- 2231296 19-04-73 US-A-3839225 01-10-74  US-A-4152301 01-05-79 AU-B- 524724 30-09-82 AU-B- 3803978 17-01-80 CA-A- 1098105 24-03-81 DE-A- 2830686 01-02-79 GB-A, B 2000985 24-01-79 JP-A- 54019494 14-02-79  US-A-4791091 13-12-88 EP-A, B 0310398 05-04-89 JP-T- 3501355 28-03-91 WO-A- 8902782 06-04-89					
SE-A- 7609423 28-02-77 US-A- 4552733 12-11-85 US-A- 4157316 05-06-79  US-A-4025606 24-05-77 GB-A- 1405405 10-09-75 DE-A- 2231296 19-04-73 US-A-3839225 01-10-74  US-A-4152301 01-05-79 AU-B- 524724 30-09-82 AU-B- 3803978 17-01-80 CA-A- 1098105 24-03-81 DE-A- 2830686 01-02-79 GB-A, B 2000985 24-01-79 JP-A- 54019494 14-02-79  US-A-4791091 13-12-88 EP-A, B 0310398 05-04-89 JP-T- 3501355 28-03-91 WO-A- 8902782 06-04-89					
US-A- 4552733 12-11-85 US-A- 4157316 05-06-79 US-A-4025606 24-05-77 GB-A- 1405405 10-09-75 DE-A- 2231296 19-04-73 US-A- 3839225 01-10-74 US-A-4152301 01-05-79 AU-B- 524724 30-09-82 AU-B- 3803978 17-01-80 CA-A- 1098105 24-03-81 DE-A- 2830686 01-02-79 GB-A, B 2000985 24-01-79 JP-A- 54019494 14-02-79 US-A-4791091 13-12-88 EP-A, B 0310398 05-04-89 JP-T- 3501355 28-03-91 WO-A- 8902782 06-04-89					
US-A- 4157316 05-06-79  US-A-4025606 24-05-77 GB-A- 1405405 10-09-75 DE-A- 2231296 19-04-73 US-A-3839225 01-10-74  US-A-4152301 01-05-79 AU-B- 524724 30-09-82 AU-B- 3803978 17-01-80 CA-A- 1098105 24-03-81 DE-A- 2830686 01-02-79 GB-A, B 2000985 24-01-79 JP-A- 54019494 14-02-79  US-A-4791091 13-12-88 EP-A, B 0310398 05-04-89 JP-T- 3501355 28-03-91 WO-A- 8902782 06-04-89					
DE-A- 2231296 19-04-73 US-A- 3839225 01-10-74  US-A-4152301 01-05-79 AU-B- 524724 30-09-82 AU-B- 3803978 17-01-80 CA-A- 1098105 24-03-81 DE-A- 2830686 01-02-79 GB-A, B 2000985 24-01-79 JP-A- 54019494 14-02-79  US-A-4791091 13-12-88 EP-A, B 0310398 05-04-89 JP-T- 3501355 28-03-91 WO-A- 8902782 06-04-89					05-06-79
DE-A- 2231296 19-04-73 US-A- 3839225 01-10-74  US-A-4152301 01-05-79 AU-B- 524724 30-09-82 AU-B- 3803978 17-01-80 CA-A- 1098105 24-03-81 DE-A- 2830686 01-02-79 GB-A, B 2000985 24-01-79 JP-A- 54019494 14-02-79  US-A-4791091 13-12-88 EP-A, B 0310398 05-04-89 JP-T- 3501355 28-03-91 WO-A- 8902782 06-04-89	115-4-4025606		GR-A- 14	 05405	10-09-75
US-A- 3839225 01-10-74  US-A-4152301 01-05-79 AU-B- 524724 30-09-82 AU-B- 3803978 17-01-80 CA-A- 1098105 24-03-81 DE-A- 2830686 01-02-79 GB-A, B 2000985 24-01-79 JP-A- 54019494 14-02-79  US-A-4791091 13-12-88 EP-A, B 0310398 05-04-89 JP-T- 3501355 28-03-91 WO-A- 8902782 06-04-89	U3-A-4023000	24 03 //			
AU-B- 3803978 17-01-80 CA-A- 1098105 24-03-81 DE-A- 2830686 01-02-79 GB-A,B 2000985 24-01-79 JP-A- 54019494 14-02-79 US-A-4791091 13-12-88 EP-A,B 0310398 05-04-89 JP-T- 3501355 28-03-91 WO-A- 8902782 06-04-89				39225	01-10-74
AU-B- 3803978 17-01-80 CA-A- 1098105 24-03-81 DE-A- 2830686 01-02-79 GB-A,B 2000985 24-01-79 JP-A- 54019494 14-02-79 US-A-4791091 13-12-88 EP-A,B 0310398 05-04-89 JP-T- 3501355 28-03-91 WO-A- 8902782 06-04-89		 01-05-79	AU-B- 5	 24724	30-09-82
US-A-4791091 13-12-88 EP-A,B 0310398 05-04-89 JP-T 3501355 28-03-91 WO-A- 8902782 06-04-89	OD W ATOCOOT	01 00 /3			
US-A-4791091 13-12-88 EP-A,B 0310398 05-04-89 JP-T- 3501355 28-03-91 WO-A- 8902782 06-04-89					
US-A-4791091 13-12-88 EP-A,B 0310398 05-04-89 JP-T- 3501355 28-03-91 WO-A- 8902782 06-04-89				30686	
US-A-4791091 13-12-88 EP-A,B 0310398 05-04-89 JP-T- 3501355 28-03-91 WO-A- 8902782 06-04-89			,		
JP-T- 3501355 28-03-91 WO-A- 8902782 06-04-89			JP-A- 540	19494	14-02-79
JP-T- 3501355 28-03-91 WO-A- 8902782 06-04-89	HS-A-4791091	13-1 <i>2</i> -88	EP-A.B 03	10398	05-04-89
WO-A- 8902782 06-04-89	03 K-4131031	13 12 00			
US-A- 4919902 24-04-90					06-04-89
			US-A- 49	19902	24-04-90